

00935290.030402

1/32

FIG. 1A

Input file Fbh56919FL2.seq
Sequence length 3003

```
TTCGGCACCAGGCTGCTGCGGGGACCTCTTTCTGAGTTACTGTGGGACCCAAAGTCTGTACGCCCTCTGCGCGTGC
AAACAGGCCACCCAGGAGAACCCAGACCTTGTCTTATTCACCCACAGCCTGGGACTGTCTTCTCCAGAGTCTCCATCAGCTT
TGCTAATCGACTGATTTGGAAATATTCCTCAACACCCACCAAGTCAAGGATACAGGCAGCAGCGGCTCCCTGTTGTAT
GGACATTTGCAACCCGAACTGATAGCTGAGTCCCTGAAAGTTTATGTTATGAAACAGAGAACTTTCATCCAGCACAT
13      M D E S A L T L G T I D V
39      GATTTGGGAATTACACTTTGTGAC ATG GAT GAA TCT GCA CTG ACC CTT GGT ACA ATA GAT GTT
33      S Y L P H S S E Y S V G R C K H T S E E
99      TCT TAT CTG CCA CAT TCA TCA GAA TAC AGT GTT GGT CGA TGT AAG CAC ACA AGT GAG GAA
53      W G E C G F R P T V F R S A T L K W K E
159     TGG GGT GAG TGT GGC TTT AGA CCC ACC GTC TTC AGA TCT GCA ACT TTA AAA TGG AAA GAA
73      S L M S R K R P P F V G R C C Y S C T P Q
219     AGC CTA ATG AGT CGG AAA AGG CCA TTT GTT GGA AGA TGT TGT TAC TCC TGC ACT CCC CAG
93      S W D K F F N P S I P S L G L R N V I Y
279     AGC TGG GAC AAA TTT TTC AAC CCC AGT ATC CCG TCT TTG GGT TTG CCG AAT GTT ATT TAT
113     I N E T H T R R H R G W L A R R L S Y V L
339     ATC AAT GAA ACT CAC ACA AGA CAC CGC GGA TGG CTT GCA AGA CGC CTT TCT TAC GTT CTT
133     F I Q E R D V H K G M F A T N V T E N V
399     TTT ATT CAA GAG CGA GAT CTG CAT ARG GGC ATG TTT GCC ACC AAT GTG ACT GAA ANT GTG
```

2/32

FIG. 1B

153 L N S S R V Q E A I A E V A A E L N P D
 459 CTG AAC AGC AGT AGA GTA CAA GAG GCA ATT GCA GAA GTG GCT GCT GAA TTA AAC CCT GAT
 173 G S A Q Q Q S K A V N K V K K K A K R I
 519 GGT TCT GCC CAG CAG CAA TCA AAA GCC GTT AAC AAA GTG AAA AAG AAA GCT AAA AGG ATT
 193 L Q E M V A T V S P A M I R L T G W V L
 579 CTT CAA GAA ATG GTT GCC ACT GTC TCA CCG GCA ATG ATC AGA CTG ACT GGG TGG GTG CTG
 213 L K L F N S F F W N I Q I H K G Q L E M
 639 CTA AAA CTG TTC AAC AGC TTC TTT TGG AAC ATT CAA ATT CAC AAA GGT CAA CTT GAG ATG
 233 V K A A T E T N L P L L F L P V H R S H
 699 GTT AAA GCT GCA ACT GAG ACG AAT TTG CCG CTT CTG TTT CTA CCA GTT CAT AGA TCC CAT
 253 I D Y L L L L T F I L F C H N I K A P Y I
 759 ATT GAC TAT CTG CTG CTC ACT TTC ATT CTC TTC TGC CAT AAC ATC AAA GCA CCA TAC ATT
 273 A S G N N L N I P I F S T L I H K L G G
 819 GCT TCA GGC AAT AAT CTC AAC ATC CCA ATC TTC AGT ACC TTG ATC CAT AAG CTT GGG GGC
 293 F F I R R R L D E T P D G R K D V L Y R
 879 TTC TTC ATA CGA CGA AGG CTC GAT GAA ACA CCA GAT GGA CGG AAA GAT GTT CTC TAT AGA
 313 A L L H G H I V E L L R Q Q Q F L E I F
 939 GCT TTG CTC CAT GGG CAT ATA GTT GAA TTA CTT CGA CAG CAG CAA TTC TTG GAG ATC TTC
 333 L E G T R S R S G K T S C A R A G L L S
 999 CTG GAA GGC ACA CGT TCT AGG AGT GGA AAA ACC TCT TGT GCT CGG GCA GGA CTT TTG TCA

3/32

FIG. 1C

V V V D T L S T N V I P D I L I I P V G 353
 GTT GTG GTA GAT ACT CTG TCT ACC AAT GTC ATC CCA GAC ATC TTG ATA ATA CCT GTT GGA 1059
 I S Y D R I I E G H Y N G E Q L G K P K 373
 ATC TCC TAT GAT CGC ATT ATC GAA GGT CAC TAC AAT GGT GAA CAA CTG GGC AAA CCT AAG 1119
 K N E S L W S V A R G V I R M L R K N Y 393
 AAG AAT GAG AGC CTG TGG AGT GTA GCA AGA GGT GTT ATT AGA ATG TTA CGA AAA AAC TAT 1179
 G C V R V D F A Q P F S L K E Y L E S Q 413
 GGT TGT GTC CGA GTG GAT TTT GCA CAG CCA TTT TCC TTA AAG GAA TAT TTA GAA AGC CAA 1239
 S Q K P V S A L L S L E Q A L L P A I L 433
 AGT CAG AAA CCG GTG TCT GCT CTA CTT TCC CTG CAG CAA GCG TTG TTA CCA GCT ATA CTT 1299
 P S R P S D A A D E G R D T S I N E S R 453
 CCT TCA AGA CCC AGT GAT GCT GCT GAT GAA GGT AGA GAC ACG TCC ATT AAT GAG TCC AGA 1359
 N A T D E S L R R R L I A N L A E H I L 473
 AAT GCA ACA GAT GAA TCC CTA CGA AGG AGG TTG ATT GCA AAT CTG GCT GAG CAT ATT CTA 1419
 F T A S K S C A I M S T H I V A C L L L 493
 TTC ACT GCT AGC AAG TCC TGT GCC ATT ATG TCC ACA CAC AAT GTG GCT TGC CTG CTC CTC 1479
 Y R H R Q G I D L S T L V E D F F V M K 513
 TAC AGA CAC AGG CAG GGA ATT GAT CTC TCC ACA TTG GTC GAA GAC TTC TTT GTG ATG AAA 1539
 E E V L A R D F D L G F S G N S E D V V 533
 GAG GAA GTC CTG GCT GAT TTT GAC CTG GGG TTC TCA GGA AAT TCA GAA GAT GTA GTA 1599

4/32

FIG. 1D

M H A I Q L L L G N C V T I T H T S R N D 553
 ATG CAT GCC ATA CAG CTG CTG GGA AAT TGT GTC ACA ATC ACC CAC ACT AGC AGG AAC GAT 1659

 E F I T P S T T V P S V F E L N F Y S 573
 GAG TTT TTT ATC ACC CCC AGC ACA ACT GTC CCA TCA GTC TTC GAA CTC AAC TTC TAC AGC 1719

 N G V L H V F I M E A I I A C S L Y A V 593
 AAT GGG GTA CTT CAT GTC TTT ATC ATG GAG GCC ATC ATA GCT TGC AGC CTT TAT GCA GTT 1779

 L N K R G L G G P T S T P P N L I S Q E 613
 CTG AAC AAG AGG GGA CTG GGG GGT CCC ACT AGC ACC CCA CCT AAC CTG ATC AGC CAG GAG 1839

 Q L V R K A A S L C Y L L S N E G T I S 633
 CAG CTG GTG CGG AAG GCG GCC AGC CTG TGC TGC CTT CTG TCC AAT GAA GGC ACC ATC TCA 1899

 L P C Q T F Y Q V C H E T V G K F I Q Y 653
 CTG CCT TGC CAG ACA TTT TAC CAA GTC TGC TGC CAT GAA ACA GTA GGA AAG TTT ATC CAG TAT 1959

 G I L T V A E H D D Q E D I E P S L A E 673
 GGC ATT CTT ACA GTG GCA GAG CAC GAT GAC GAT GAC CAG GAA GAT ATC AGT CCT AGT CTT GCT GAG 2019

 Q Q W D K K L P E P L S W R S D E E D E 693
 CAG CAG TGG GAC AAG AAG CTT CCA GAA CCT TTG TGT TGG AGA AGT GAT GAA GAA GAT GAA 2079

 D S D F G E E Q R D C Y L X V S Q S K E 713
 GAC AGT GAC TTT GGG GAG GAA CAG CGA GAT TGC TAC CTG AAG GTG AGC CAA TCC AAG GAG 2139

 H Q Q Q F I T F L Q R L L G P L L E A Y S 733
 CAC CAG CAG TTT ATC ACC TTC TTA CAG AGA CTC CTT GGG CCT TTG CTG GAG GCC TAC AGC 2199

May-17-02 09:30pm

From-LAH1 ROCKFIELD, LLP

6177424214

T-652 P.27/54 F-983

5/32

FIG. 1E

S A A I F V H N F S G P V P E P E Y L Q 753
 TCT GCT GCC ATC TTT GTT CAC AAC TTC AGT GGT CCT GTT CCA GAA CCT GAG TAT CTG CAA 2259

 K L H K Y L I T R T E R N V A V Y A E S 773
 AAG TTG CAC AAA TAC CTA ATA ACC AGA ACA GAA AAT GTT GCA GTA TAT GCT GAG AGT 2319

 A T Y C L V K N A V K M F K D I G V F K 793
 GCC ACA TAT TGT CTT GTG AAG AAT GCT GTG AAA ATG TTT AAG GAT ATT GGG GTT TTC AAG 2379

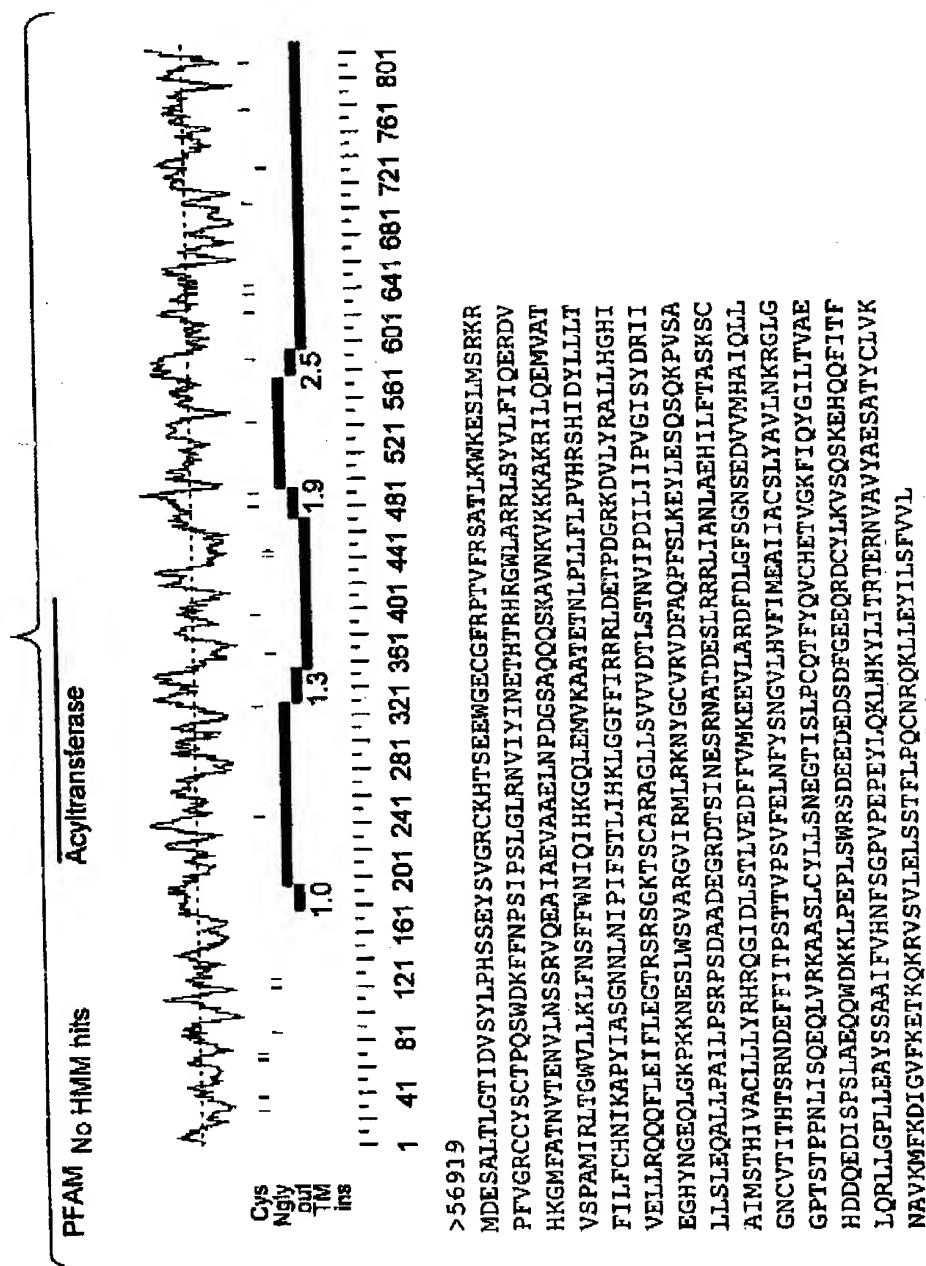
 E T K Q K R V S V L E L S S T F L P Q C 813
 GAG ACC AAA CAA AAG AGA GTG TCT GTT TTA GAA CTG AGC AGC ACT TTT CTA CCT CAA TGC 2439

 N R Q K L L E Y I L S F V V L * 829
 AAC CGA CAA AAA CTT CTA GAA TAT ATT CTG AGT TTT GTG GTG CTG TAG 2487

 GTAACGTGTGGCACTGCGCAATGAAGGTCATGAGATGAGTTCCCTTGTAGGTACCAGCTTCTGGCTCAAGAGTTTGA
 AGGTGCCCTTCCGAGGSETCAGGCCCTGCCCTGTGTCGAGTATCTCCTGGAGACAAAGTGCCCTTCNCCCTCCATGGATC
 TGAGATCTTCCCGAGCTTT

6/32

FIG. 2



7/32

FIG. 3A

Protein Family / Domain Matches, HMMer version 2

Searching for complete domains in PFAM
 humpfam - search a single seq against HMM database
 HMMER 2.1.1 (Dec 1998)
 Copyright (C) 1992-1998 Washington University School of Medicine
 HMMER is freely distributed under the GNU General Public License (GPL).

 HMM file: /prod/ddm/seganal/PFAM/pfam6.4/Pfam
 Sequence file: /prod/ddm/wspace/orfanal/oa-script.21255.seq

Query: 56919

Scores for sequence family classification (score includes all domains):

Model	Description	Score	E-value	N
-----	-----	-----	-----	-----
Acyltransferase	Acyltransferase	126.1	6.4e-34	1

Parsed for domains:

Model	Domain	seq-f	seq-t	hmm-f	hmm-t	score	E-value
-----	-----	-----	-----	-----	-----	-----	-----
Acyltransferase	1/1	215	412 ..	1	195 []	126.1	6.4e-34

8/32

FIG. 3B

Alignments of top-scoring domains:

Acyitransferase: domain 1 of 1, from 215 to 412: score 126.1, E = 6.4e-34

*->lenlpkkgaivvsNHRsYldilvlssaalprigpwlvrilvfiakke

+ +++++ p ++ + HRS++D+1+1 ++1++++ ++ +ia ++

56919 215 KAATETNLPLLLFLPVHRSHIDYLLLTFFILFCHN-----IXAPYIASGN 257

llkvPllfGwlmlragaiifidRmra.....kdalaaadelvrvlellrk

+1++P+ f++l+ ++g +fi+R+ ++++++kd l++a+ + + tellr+

56919 258 NLNIPI-FSTLIHKLGSGFFIRRLDEtpdgrKDVLYRALLHGHIVELLNQ 306

grsvliFPEGThrsrgellppfKkGla.....afrlAlkagvpivPvviv

+ + iF EGTrsrsg++ + ++G++ + + ++ ++ i+Pv+i

56919 307 QQFLEIFLEGTrSRSGKTSC-ARAGLLsvvvdTLSTNVIPDILIIIPVGI- 354

sgteelepknasgklrlarkkgpvtvrvlppipld..pedikelaelr

s++ ++e ++++ + +++kk+++++ v +++ +++++ +

56919 355 SYDRIIEGHYNSEQ--LGKPKKNESLMSVARGVIRmlrKNYGCVRVDFAQ 402

dilvqaleel<*

+ +++++ e+

56919 403 PFSLEYLES 412

09935290.030402

9/32

FIG. 4A

ProDom Matches ProdomId	Start	End	Description	Score
View Prodom PD347660	1	55	p2001.1 (2) PLSB(2) // ACYLTRANSFERASE PHOSPHOLIPID BIOSYNTHESIS PRECURSOR TRANSMEMBRANE GLYCEROL-3-PHOSPHATE GPAT MITOCHONDRION MITOCHONDRIAL	250
View Prodom PD087501	51	158	p2001.1 (1) // AIP2-DLD1	77
View Prodom PD353751	56	152	p2001.1 (2) PLSB(2) // ACYLTRANSFERASE PHOSPHOLIPID BIOSYNTHESIS PRECURSOR TRANSMEMBRANE GLYCEROL-3-PHOSPHATE GPAT MITOCHONDRION MITOCHONDRIAL	500
View Prodom PD037846	128	259	p2001.1 (15) PLSB(5) DAPT(2) // ACYLTRANSFERASE GLYCEROL-3-PHOSPHATE PHOSPHOLIPID GPAT BIOSYNTHESIS MITOCHONDRIAL TRANSMEMBRANE PRECURSOR MITOCHONDRION	258
View Prodom PD042466	259	590	p2001.1 (16) PLSB(4) DAPT(2) // ACYLTRANSFERASE GLYCEROL-3-PHOSPHATE BIOSYNTHESIS PHOSPHOLIPID GPAT PRECURSOR MITOCHONDRIAL TRANSMEMBRANE MITOCHONDRION	413
View Prodom PD025192	462	649	p2001.1 (4) PLSB(3) // ACYLTRANSFERASE PHOSPHOLIPID MITOCHONDRIAL BIOSYNTHESIS PRECURSOR TRANSMEMBRANE GLYCEROL-3-PHOSPHATE GPAT MITOCHONDRION	

May-17-02 09:31pm

From-LAH... ROCKFIELD, LLP

6177424214

T-652 P.32/54 F-983

10/32

FIG. 4B

458

View Prodom PD042027 465 673 p2001.1 (6) PLSB(2) // ACYLTRANSFERASE
GLYCEROL-3-PHOSPHATE MEMBRANE
PHOSPHOLIPID GPAT BIOSYNTHESIS MUTANT

80

View Prodom PD042760 650 828 p2001.1 (2) PLSB(2) // ACYLTRANSFERASE
PHOSPHOLIPID BIOSYNTHESIS PRECURSOR
TRANSMEMBRANE GLYCEROL-3-PHOSPHATE
GPAT MITOCHONDRION MITOCHONDRIAL

799

View Prodom PD042760

>PD042760 p2001.1 (2) PLSB(2) // ACYLTRANSFERASE PHOSPHOLIPID BIOSYNTHESIS
PRECURSOR TRANSMEMBRANE GLYCEROL-3-PHOSPHATE GPAT MITOCHONDRION
MITOCHONDRIAL
Length = 179

Score = 799 (286.3 bits), Expect = 1.2e-79, P = 1.2e-79
Identities = 155/179 (86%), Positives = 161/179 (89%)

Query: 650 FIQYGILTVAEHDDQEDISPSLAEOQWKKLPEPLSWRXXXXXXXXXXXXXQDDCYLKVS 709
FIQYGILTVAE DDQED+SP LAEQW+KKLPEPL+WR QDDCYLKVS
Sbjct: 1 FIQYGILTVAEQDDQEDVSPGLAEQWKKLPEPLNWRSEDEDESDFGEEQDDCYLKVS 60

Query: 710 QSKHQQTITFLORLLGPLLEAYSSAAIFVHNFSGPVPEPEYLOKLHKYLITRERNVAV 769
Q+KEHQQTITFLORLLGPLLEAYSSAAIFVHNF GPVPE EYLOKLH+YLLITRERNVAV
Sbjct: 61 QAKHQQTITFLORLLGPLLEAYSSAAIFVHNFSGPVPESEYLOKLHRYLLITRERNVAV 120

Query: 770 YAESATYCLVKNVAVKMFKDGVFKETKQKRVSVLELSSTFLPQCNRQKLEYILSFVVL 828
YAESATYCLVKNVAVKMFKDGVFKETKQKRVSVLELS+TFLPQCNRQKLEYILSFVVL
Sbjct: 121 YAESATYCLVKNVAVKMFKDGVFKETKQKRVSVLELSSTFLPQCNRQKLEYILSFVVL 179

11/32

FIG. 4C

View Prodom PD353751

>PD353751 p2001.1 (2) PLSB(2) // ACYLTRANSFERASE PHOSPHOLIPID BIOSYNTHESIS
 PRECURSOR TRANSMEMBRANE GLYCEROL-3-PHOSPHATE GPAT MITOCHONDRIAL

Length = 97

Score = 500 (181.1 bits), Expect = 2.1e-47, P = 2.1e-47
 Identities = 90/97 (92%), Positives = 97/97 (100%)

Query: 56 MSRKRPVGRCCYCTPQSWDKFFNPSPSLGLRNVIYINETHTRHRGWLARRLSYVLFI 115
 MSRKRPVGRCCYCTPQSW++FFNPSPSLGLRNVIYINETHTRHRGWLARRLSY+LF+
 Sbjct: 1 MSRKRPVGRCCYCTPQSWERFFNPSPSLGLRNVIYINETHTRHRGWLARRLSYILFV 60

Query: 116 QERDVHKGMFATNVNTENVLNSSRVQEAIAEVAALNP 152
 QERDVHKGMFAT++T+NVLNSSRVQEAIAEVAALNP
 Sbjct: 61 QERDVHKGMFATSI TDNVLNSSRVQEAIAEVAALNP 97

View Prodom PD025192

>PD025192 p2001.1 (4) PLSB(3) // ACYLTRANSFERASE PHOSPHOLIPID MITOCHONDRIAL
 BIOSYNTHESIS PRECURSOR TRANSMEMBRANE GLYCEROL-3-PHOSPHATE GPAT
 MITOCHONDRIAL

Length = 194

Score = 458 (166.3 bits), Expect = 8.7e-43, P = 8.7e-43
 Identities = 95/192 (49%), Positives = 126/192 (65%)

Query: 462 RRLIANLAEHILFTASKSCAISMTHIVACILLYRHRQGLDLSTLVEDFFVMKEVLARDF 521
 R LI ++ EH++F S C+INSTH+VACILL R R G+ STL ED + E+LA
 Sbjct: 3 RNLIRSIGEHVVFDSCMMCSINSTHVACILLTRWRNGVHRSTLEEDCDWLCEKILAEAG 62

09935290.030402

12/32

FIG. 4D

Query: 522 DL-GFSGNS---EDVVMHAIQLLNCVITHTSRNDEFFITPSTTVPSVFELNFYSNGVL 577
 D+ GFSG S +V +A +LLG+CVT+T RNDFF+I+P +VPS EL +YSN V+

Sbjct: 63 DIVGFSKSTKGSQIVKYACELLGSCVTVTDEDRNDEFFIYSPKNSVPSEFIELAYYSNSVI 122

Query: 578 HVFIMEAIIACSLYAVLNKRGGLGPTSTPPNLLSQEQIVRKAASLCYLLSNEGTSISLPCQ 637
 F +++IIAC++Y++ NK GG NLISQEQLV A SLC L E PCQ

Sbjct: 123 CHFALKSIIACTIYSLPNKTKNGGAGLGNLISQEQIVEDALSCLDWLQYEFMFRCPCQ 182

Query: 638 TFYQVCHETVGK 649
 T ++CH T+GK

Sbjct: 183 TLRELCHNTLGK 194

View Prodom PD042466

>PD042466 p2001.1 (16) PLSB(4) DAPT(2) // ACYLTRANSFERASE GLYCEROL-3-PHOSPHATE
 BIOSYNTHESIS PHOSPHOLIPID GPAT PRECURSOR MITOCHONDRIAL TRANSMEMBRANE
 MITOCHONDRION
 Length = 299

Score = 413 (150.4 bits), Expect = 4.7e-42, Sum P(2) = 4.7e-42
 Identities = 79/152 (51%), Positives = 108/152 (71%)

Query: 259 LNIPIFSTLIHKLGGFFIRRLDETDPGRKDVLYRALLHGHIVELLRQQQFLEIFLECTR 318
 L++PI +L+ + G FFIRR D TP+G+ D LYRA+ H ++ +L+ + +E F+EGTR

Sbjct: 2 LSPIMGSLLRRTGAFFIRRSFDPTEGKGDQLYRAVHEYVAQLISKGYNIEFFIEGTR 61

Query: 319 SRSGKTSCARAGLLSVVVDTLSTNVIPDILLIPVGISYDRIIEGH-YNGEQLGKPKKNES 377
 SR+GK + GLLS+VV+ +PDIL++PV ISYDRIIEG+ Y E G PKK ES

Sbjct: 62 SRTGKMLPPKGTGLLSMVVVEAFLRGSVPDILLVPVVISYDRIIEGNTYAHELARGAPKKES 121

13/32

FIG. 4E

Query: 378 LWSVARGVIRMLRKKNYGCVRVDFEAQPFSLKEY 409

LW + RGV +ML++NYG V VDF +P SL+EY

Sbjct: 122 LWQLFRGVRKMLKRNNGOVYVDFGEPISLREY 153

Score = 64 (27.6 bits), Expect = 4.7e-42, Sum P(2) = 4.7e-42
 Identities = 30/143 (20%), Positives = 59/143 (41%)

Query: 453 RNATDESILRRRLIANLAEHILFTASKSCAIMSTHIVACLLLYRHRQIGIDLSTL---VEDF 509
 RN ++ + +R + ++ + + ++ + +T +V+ LLL + L L ++D

Sbjct: 160 RNTYNCAPKRLALQKMSFEVAVRWIRILOATPVTATGLYSALLLTTRGTALTLDQLHHTLQDS 219

Query: 510 FVMKEEVLARDFDLGFSGNSEDDVVMHAIQLL---GNCVTITHTSRNDFFITPTTVPVSF 567
 E + S + V A L G+ VT + R ++I P + F

Sbjct: 220 LDYLERKQSPVSTSAIRLRSREGVRAAADALSNGHPVTRVDSGREPVWYIAPDDEHAAAF 279

Query: 568 ELNFYSNGVLHVVFIMEAIIACSL 590

Y N V+H F+ +I+ +L

Sbjct: 280 -----YRNSVIHAFLETTSIVELAL 298

View Prodom PD037B46

>PD037B46 p2001.1 (15) PLSH(5) DAPT(2) // ACYLTRANSFERASE GLYCEROL-3-PHOSPHATE
 PHOSPHOLIPID GPAT BIOSYNTHESIS MITOCHONDRIAL TRANSMEMBRANE PRECURSOR
 MITOCHONDRION
 Length = 345

Score = 258 (95.9 bits), Expect = 3.4e-21, P = 3.4e-21
 Identities = 56/132 (42%), Positives = 79/132 (59%)

14/32

FIG. 4F

Query: 128 NVTENVLNSSRVQEAIAEVAELNPDGSAQQQXXXXXXXXXXXXXRILOEMVATVSPAMIR 187
 N+ +NVLNS + I + A++ S IL EM T++ MIR
 Sbjct: 223 NLKKNVLNSEEIHVVIEQ-----EAKESSTSIDKVRREAREILDEMSHTLNMGMIR 273

Query: 188 LTGMVLLKLFNSFFWNIIQIHKGQLEMVKAATETNPLFLFPVHRSHIDYLLLTFFILFCHN 247
 GWVL K+FN F I +++ Q+E +K ATE P+++LP HRSHIDYLL+FIL+ ++
 Sbjct: 274 FCGWVLSKIFNRIFSGICVNEEQIEKIKRATEQGHPIYLPVHRSHIDYLLLTSEILYHYD 333

Query: 248 IKAPYIASGNL 259
 IK P+IA+G NL
 Sbjct: 334 IKVPHIAAGMNL 345

View Prodom PD347660

>PD347660 p2001.1 (2) PLSB(2) // ACYLTRANSFERASE PHOSPHOLIPID BIOSYNTHESIS
 PRECURSOR TRANSMEMBRANE GLYCEROL-3-PHOSPHATE GPAT MITOCHONDRION
 MITOCHONDRIAL
 Length = 55

Score = 250 (93.1 bits), Expect = 2.4e-20, F = 2.4e-20
 Identities = 43/55 (78%), Positives = 53/55 (96%)

Query: 1 MDESALTIGTIDVSYLPHSSEYSVGRCKHISEWGECCGFRPTVFRSATLKKWESL 55
 M+ES++T+GTIDVSYLP+SSEYS+GRCKHT+E+W +CGF+PT FRSATLKKWESL
 Sbjct: 1 MEESSTVIGTIDVSYLPNSSEYSIGRCKHTNEDWVDCGKPTFFRSATLKKWESL 55

09935290.030402

15/32

FIG. 4G

View Prodrom PD042027

>PD042027 p2001.1 (6) PLSB(2) // ACYLTRANSFERASE GLYCEROL-3-PHOSPHATE MEMBRANE
 PHOSPHOLIPID GPAT BIOSYNTHESIS MUTANT
 Length = 345

Score = 80 (33.2 bits), Expect = 0.11, Sum P(2) = 0.10
 Identities = 31/129 (24%), Positives = 60/129 (46%)

Query: 465 IANLAEHILFTASKSKAIMSTHIVACILLYRHRQIGIDLSTLVE--DFFVMKEEVLARDFD 522
 + +LA+ I+ + + A+ ++ A LL ++ + L+E D ++ + D

Sbjct: 32 VNHLAKQIMTHINDAAAANPMLCATALLSTRQALGEEQLIEQLDCYLKLLRNVPYSTD 91

Query: 523 LGFSGNS-EDVVMHAIQ--LLGNCVTITHTSRNDEFFITPTTVPSPVFELNFYSNGVLHV 579
 ++ E ++ HA Q LLG VT+ + D + V + +Y N VILH+

Sbjct: 92 ATLPDHTPERLIEHAEQMNLLG--VTVEKDTLGLDLRLDRDNAVL----MTYRNNVNLHL 145

Query: 580 FIMEAIIAC 588
 F + A++AC

Sbjct: 146 FALPALVAC 154

Score = 62 (26.9 bits), Expect = 0.11, Sum P(2) = 0.10
 Identities = 22/105 (20%), Positives = 48/105 (45%)

Query: 569 LNFYSNGVLHVFIIMEAIIACSLYAVLNKRGGLGGPTSTPPNLI SQEQLVRKAASLCYLLSN 628
 + +Y N VLH+F + A++AC N+R IS++ I+R +L L

Sbjct: 135 MTYRNNVNLHLFALPALVACCFKN--NRR-----ISRALLRFVRALYPFLQA 180

Query: 629 EGTISLPCQTFYQVCHETVCKFIQYGLTVAEHDDQEDISPSLAE 673
 E + + + +F++ G+L A + + + + + +

Sbjct: 181 ELFLRWNEDELNDHIDQWINEFVRQGLLSAGNQEDDTLFRNTSQ 225

09935290.030402

16/32

FIG. 4H

View Prodom PD087501

>PD087501 p2001.1 (1) // AIP2-DLD1
 Length = 170

Score = 77 (32.2 bits), Expect = 5.1, P = 0.994
 Identities = 31/114 (27%), Positives = 44/114 (38%)

Query: 51 WKES--LMSRKRPFVGRCCYCTPQSWDKFFNPSLGLRNVIIYNETHTRHGWLARR 108
 W ES L+ RK F RCC F K + L N +H W
 Sbjct: 12 WNESEVLVDRKSKFQARCC----PLQNKDIPSLQELTONNKSVSKASHMHMYAWRTAE 67

Query: 109 LSYVLFQERDVHKGMFATNVTVTENVLNSR---VQ-EAIAEVAELNPDGSAQQ 158
 +S L+Q+ KG A + +N SR VQ + I + A+ G+ Q+
 Sbjct: 68 VSNNLHLQOEQKKKGNKANKSNNSHVNKSRNITVQPKNIEQGCADCGEAGAGOR 121

17/32

FIG. 5A

	10	20	30	40	
1	MDESALTGLTIDVSYLPHSSEYSVGRCKHTSEEWGECGFR				56919.pro
1	MEESSVTVGTIDVSYLPSSEYSLGRCKHTSEDWVDCGFK				MouseGPAT.PRO
1	MEESSVTIGTIDVSYLPNSSEYSLGRCKHTNEDWVDCGFK				RatGPAT.PRO
	50	60	70	80	
41	PTVFRSATLKWKESLMSRKRPFVGRCCYSCTPQSWDKFFN				56919.pro
41	PTFFRSATLKWKESLMSRKRPFVGRCCYSCTPQSWERFFN				MouseGPAT.PRO
41	PTFFRSATLKWKESLMSRKRPFVGRCCYSCTPQSWERFFN				RatGPAT.PRO
	90	100	110	120	
81	PSIPSLGLRNVIIYNETHTRHRGWLARRLSYVLFQERDV				56919.pro
81	PSIPSLGLRNVIIYNETHTRHRGWLARRLSYILFVQERDV				MouseGPAT.PRO
81	PSIPSLGLRNVIIYNETHTRHRGWLARRLSYILFVQERDV				RatGPAT.PRO
	130	140	150	160	
121	HKGMFATNVTVENVLNSSRVQEAIAEVAELNPDGSAQQQS				56919.pro
121	HKGMFATSVTVENVLNSSRVQEAIAEVAELNPDGSAQQQS				MouseGPAT.PRO
121	HKGMFATSITDNVLNSSRVQEAIAEVAELNPDGSAQQQS				RatGPAT.PRO
	170	180	190	200	
161	KAVNKVKKKAKRILQEMVATVSPAMIRLTGWVLLKLFNSF				56919.pro
161	KAIQKVKKRARKILQEMVATVSPGIRLTGWVLLKLFNSF				MouseGPAT.PRO
161	KAIQKVKKRARKILQEMVATVSPGIRLTGWVLLKLFNSF				RatGPAT.PRO
	210	220	230	240	
201	FWNIQIHKGQLEMVKAATETNPLFLPVHRSHIDYLLLT				56919.pro
201	FWNIQIHKGQLEMVKAATETNPLFLPVHRSHIDYLLLT				MouseGPAT.PRO
201	FWNIQIHKGQLEMVKAATETNPLFLPVHRSHIDYLLLT				RatGPAT.PRO
	250	260	270	280	
241	FILFCHNIKAPYIASGNLNIPIFSTLIHKLGGFFIRRL				56919.pro
241	FILFCHNIKAPYIASGNLNIPIVSTLIHKLGGFFIRRL				MouseGPAT.PRO
241	FILFCHNIKAPYIASGNLNIPIFSTLIHKLGGFFIRRL				RatGPAT.PRO
	290	300	310	320	
281	DETPDGRKDVLYRALLHGHIVELLRQQQFLEIFLEGTRSR				56919.pro
281	DETPDGRKDILYRALLHGHVVELLRQQQFLEIFLEGTRSR				MouseGPAT.PRO
281	DETPDGRKDILYRALLHGHIVELLRQQQFLEIFLEGTRSR				RatGPAT.PRO
	330	340	350	360	
321	SGKTSCARAGLLSVVVDTLSTNVIPDILIPVGISYDRII				56919.pro
321	SGKTSCARAGVLSVVNTLSSNTIPDILIPVGISYDRII				MouseGPAT.PRO
321	SGKTSCARAGLLSVVVDTLSSNTIPDILIPVGISYDRII				RatGPAT.PRO

18/32

FIG. 5B

	370	380	390	400	
361	EGHYNGEQLGKPKKNESLWSVARGVIRMLRKNYGCVRVDF				56919.pro
361	EGHYNGEQLGKPKKNESLWSVARGVIRMLRKNYGYVRVDF				MouseGPAT.PRO
361	EGHYNGEQLGKPKKNESLWSVARGVIRMLRKNYGYVRVDF				RatGPAT.PRO
	410	420	430	440	
401	AQPFSLKEYLESQSQKPVSAALLSLEQALLPAILPSRPSDA				56919.pro
401	AQPFSLKEYLEGQSQKPVSAPLSLEQALLPAILPSRPNDV				MouseGPAT.PRO
401	AQPFSLKEYLEGQSQKPVSAPLSLEQALLPAILPSRPDAA				RatGPAT.PRO
	450	460	470	480	
441	ADEGRDTSINESRNATDESLRRRLIANLAEHILFTASKSC				56919.pro
441	ADEHQDLSINESRNPADAFRRRLIANLAEHILFTASKSC				MouseGPAT.PRO
441	AAEHEDMSINESRNAADEAFRRRLIANLAEHILFTASKSC				RatGPAT.PRO
	490	500	510	520	
481	AIMSTHIVACLLLYRHRQIDLSTLVEDFFVMKEEVLARD				56919.pro
481	AIMSTHIVACLLLYRHRQGIHLSTLVEDFFVMKEEVLARD				MouseGPAT.PRO
481	AIMSTHIVACLLLYRHRQGIHLSTLVEDFFVMKEEVLARD				RatGPAT.PRO
	530	540	550	560	
521	FDLGFSGNSEDEVVMHAIQLLGNCVTITHTSRNDEFFITPS				56919.pro
521	FDLGFSGNSEDEVVMHAIQLLGNCVTITHTSRKDEFFITPS				MouseGPAT.PRO
521	FDLGFSGNSEDEVVMHAIQLLGNCVTITHTSRKDEFFITPS				RatGPAT.PRO
	570	580	590	600	
561	TTVPSVFELNFYSNGVLHVFIMEAIIACSLYAVLNKRGLG				56919.pro
561	TTVPSVFELNFYSNGVLHVFIMEAIIACSLYAVLNKRCSG				MouseGPAT.PRO
561	TTVPSVFELNFYSNGVLHVFIMEAIIACSLYAVQNKRGSG				RatGPAT.PRO
	610	620	630	640	
601	GPTSTPPNLISQEQLVRKAASLCYLLSNEGTTISLPCQTFY				56919.pro
601	GPTSTPPNLISQEQLVRKAASLCYLLSNEGTTISLPCQTFY				MouseGPAT.PRO
601	GPTSTPPNLISQEQLVRKAASLCYLLSNEGTTISLPCQTFY				RatGPAT.PRO
	650	660	670	680	
641	QVCHETVGKFIQYGILTVAEHDDQEDISPGLAEQQWDKKL				56919.pro
641	QVCHETVGKFIQYGILTVAEQDDQEDVSPGLAEQQWDKKL				MouseGPAT.PRO
641	QVCQETVGKFIQYGILTVAEQDDQEDVSPGLAEQQWNKKL				RatGPAT.PRO
	690	700	710	720	
681	PEPLSWRSDEEDEDSDFGEEQRDCYLKVSQSKEHQOFITF				56919.pro
681	PE-LNWRSD E E D E D S D F G E E Q R D C Y L K V S Q S K E H Q Q F I T F				MouseGPAT.PRO
681	PEPLNWRSD E E D E D S D F G E E Q R D C Y L K V S Q A K E H Q Q F I T F				RatGPAT.PRO

May-17-02 09:34pm From-LAH1 ROCKFIELD, LLP

6177424214

T-652 P.41/54 F-983

19/32

FIG. 5C

	730	740	750	760	
721	LQRLLGPLLEAYSSAAIFVHNFSGPVPEPEY LQKLHKYLI				56919.pro
721	LQRLLGPLLEAYSSAAIFVHNFSGPVPEPEY LQRLHKYLI				MouseGPAT.PRO
721	LQRLLGPLLEAYSSAAIFVHTFRGPVPEPEY LQRLHKYLI				RatGPAT.PRO
	770	780	790	800	
761	TRTERNVAVYAESATYCLVKNVAVKMFKDIGVFKETKQKRV				56919.pro
761	TRTERNVAVYAESATYCLVKNVAVKMFKDIGVFKETKQKRV				MouseGPAT.PRO
761	TRTERNVAVYAESATYCLVKNVAVKMFKDIGVFKETKQKRA				RatGPAT.PRO
	810	820			
801	SVLELSSTFLPQCNRQKLLEYILSFVVL				56919.pro
801	SVLELSSTFLPQCNRQKLLEYILSFVVL				MouseGPAT.PRO
801	SVLELSSTFLPQCNRQKLLEYILSFVVL				RatGPAT.PRO

May-17-02 09:35pm

From-LAH1 ROCKFIELD, LLP

6177424214

T-652 P.42/54 F-983

20/32

FIG.6

Acyltransferase catalytic motif-I

IFLEGTRSR	56919.pro
IFLEGTRSR	MouseGPAT.PRO
IFLEGTRSR	RatGPAT.PRO
YFVEGGRSR	EcoliGPAT.PRO

Acyltransferase catalytic motif-II

HRSHID	56919.pro
HRSHID	MouseGPAT.PRO
HRSHID	RatGPAT.PRO
HRSHMD	EcoliGPAT.PRO

Acyltransferase catalytic motif-III

ILIIPV	56919.pro
ILVIPV	MouseGPAT.PRO
ILVIPV	RatGPAT.PRO
ITLIPI	EcoliGPAT.PRO

Acyltransferase signature motif

GGFFIRR	56919.pro
GGFFIRR	MouseGPAT.PRO
GGFFIRR	RatGPAT.PRO
GAFFIRR	EcoliGPAT.PRO

May-17-02 09:35pm

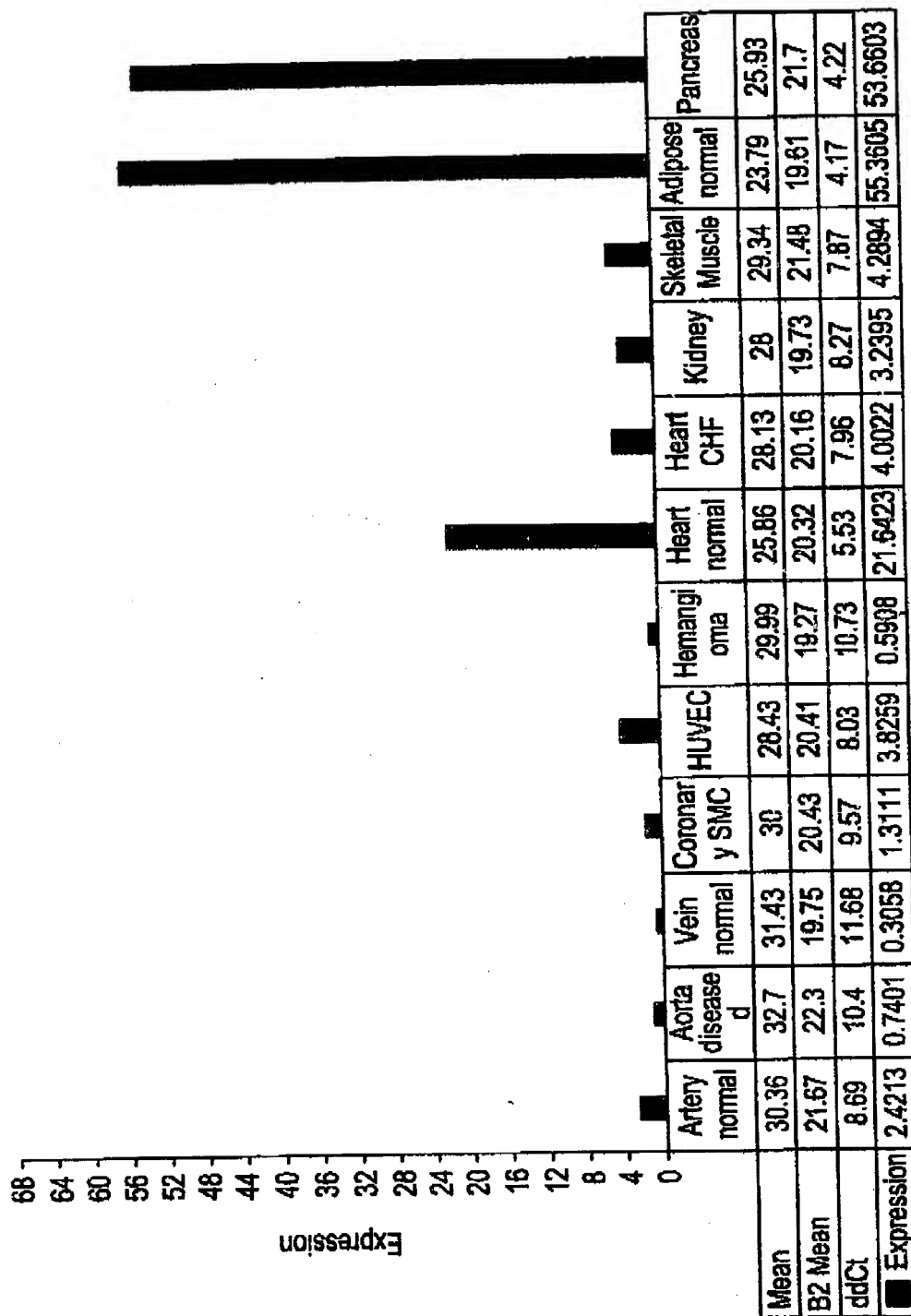
From-LAH1 COCKFIELD, LLP

6177424214

T-652 P.43/54 F-983

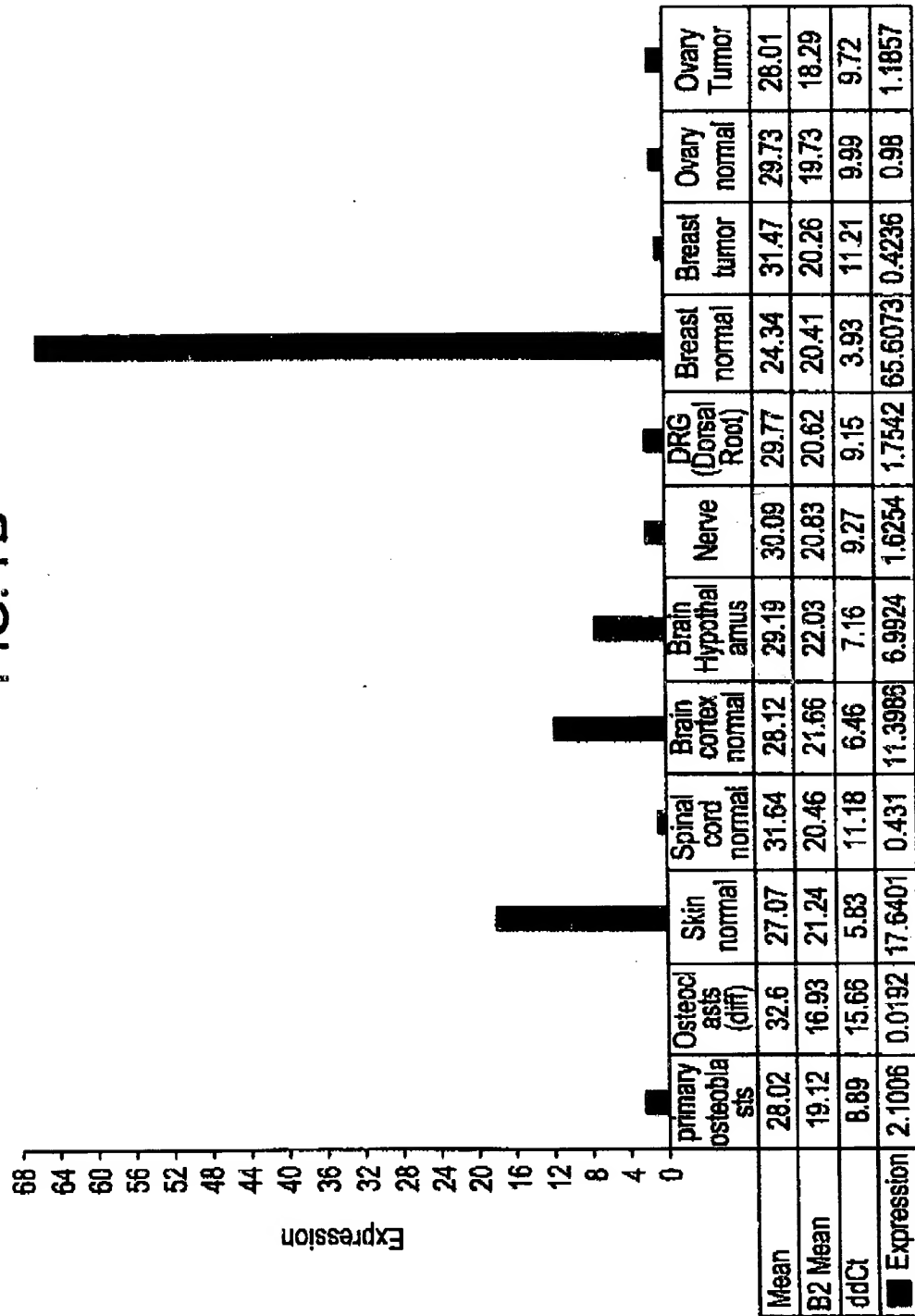
21/32

FIG. 7A



22/32

FIG. 7B



May-17-02 09:35pm

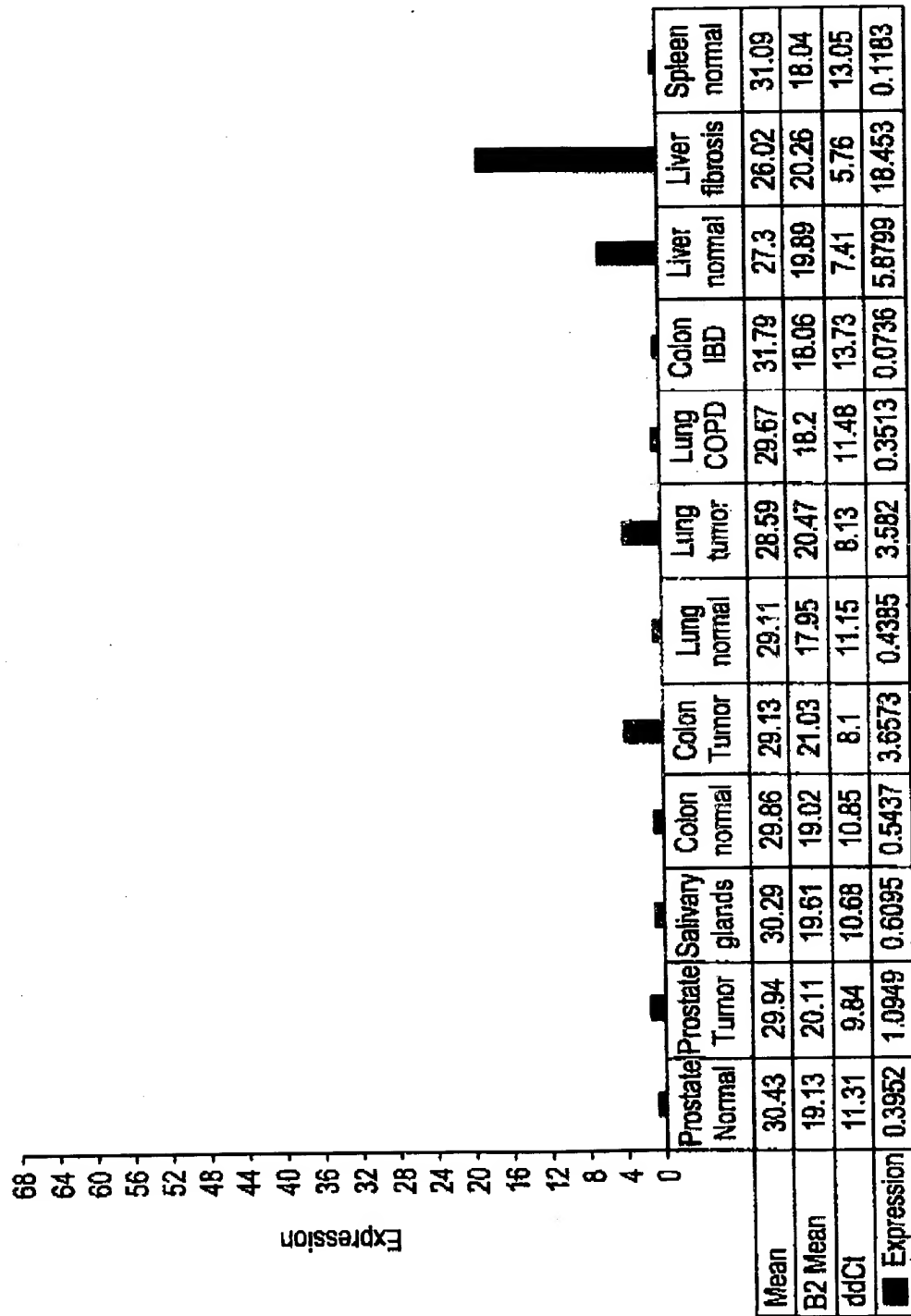
From-LAH1 ROCKFIELD, LLP

6177424214

T-652 P.45/54 F-983

23/32

FIG. 7C



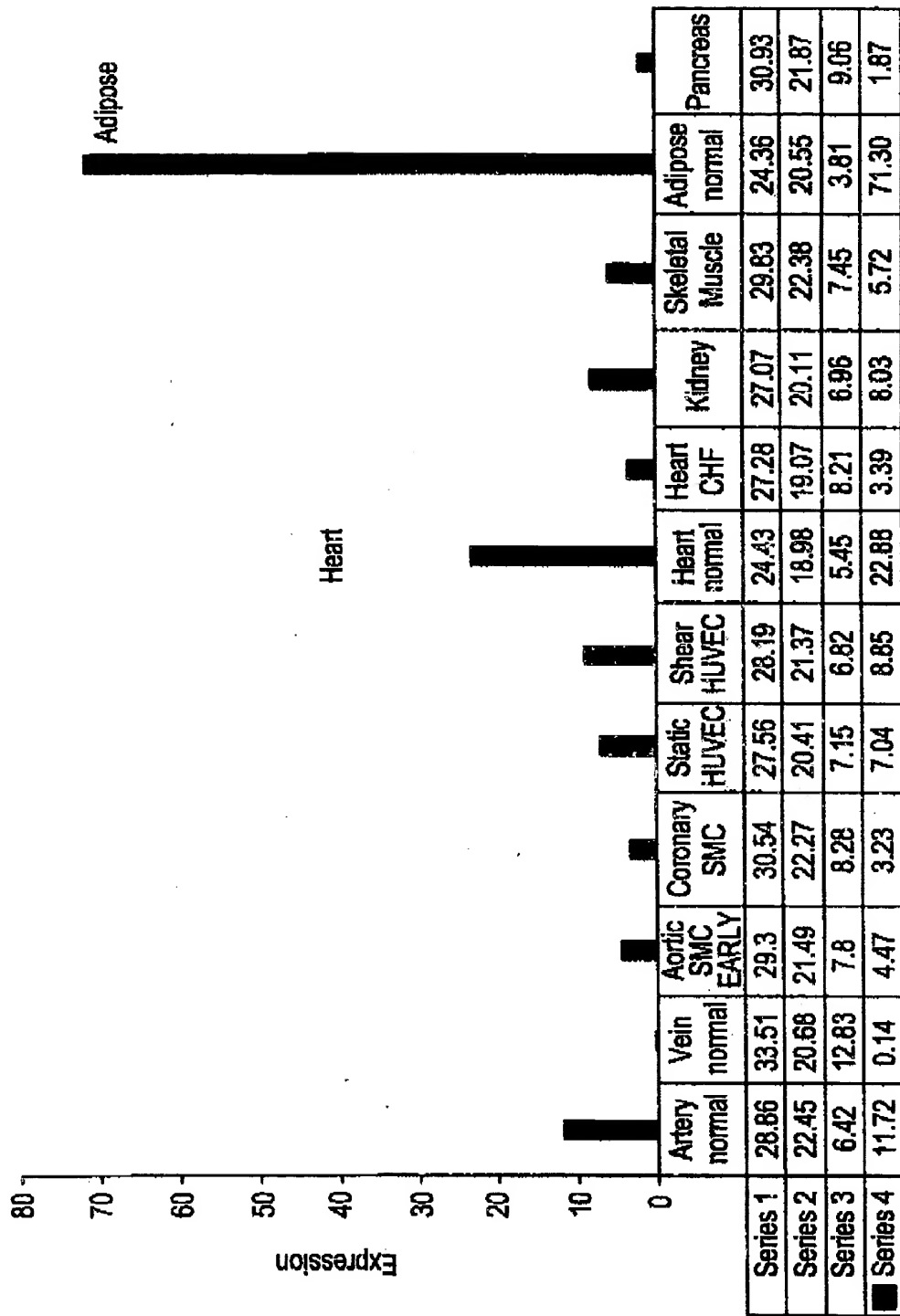
24/32

FIG. 7D

Expression	Tonsil normal	Lymph node normal	Small intestine normal	Macroph ages	Synoviu m	BM- MNC	Activate d PBMC	Neutrop hils	Megakar yocytes	Erythro id	positive control
Mean	29.32	31.91	31.66	34.2	33.57	34.91	31.78	30.61	30.13	28.29	28.32
B2 Mean	18.11	19.79	19.7	16.75	18.7	18.13	17.61	17.54	18.28	20.27	20.84
ddCt	11.21	12.13	11.97	17.45	14.87	16.77	14.16	13.07	11.85	8.02	7.48
Expression	0.4236	0.2239	0.2493	0.0056	0.0334	0.0089	0.0544	0.1163	0.2709	3.8525	5.6014

25/32

FIG. 8A



May-17-02 09:36pm

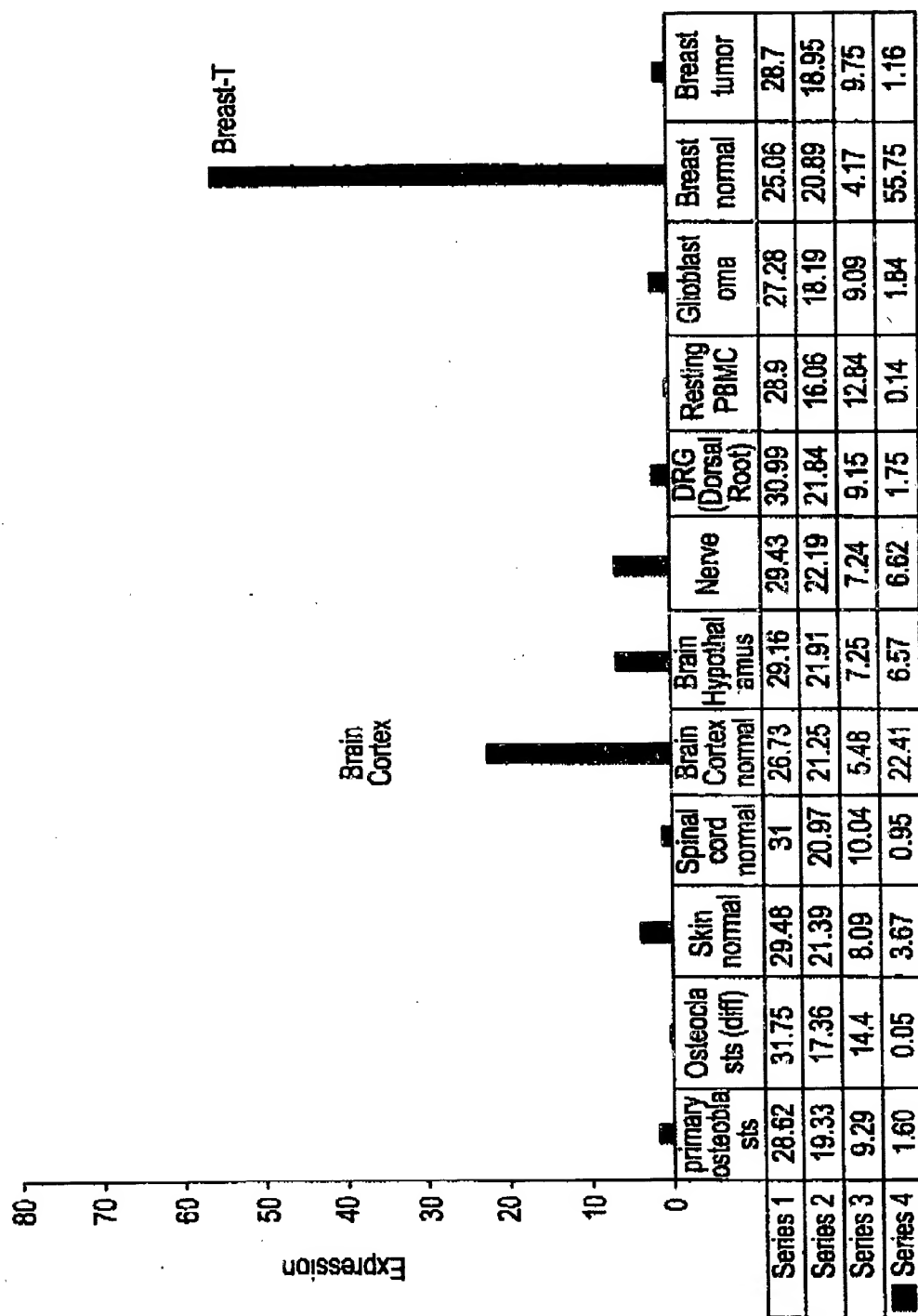
From-LAH LOCKFIELD, LLP

6177424214

T-652 P.48/54 F-983

26/32

FIG. 8B



May-17-02 09:36pm

From-LAH COCKFIELD, LLP

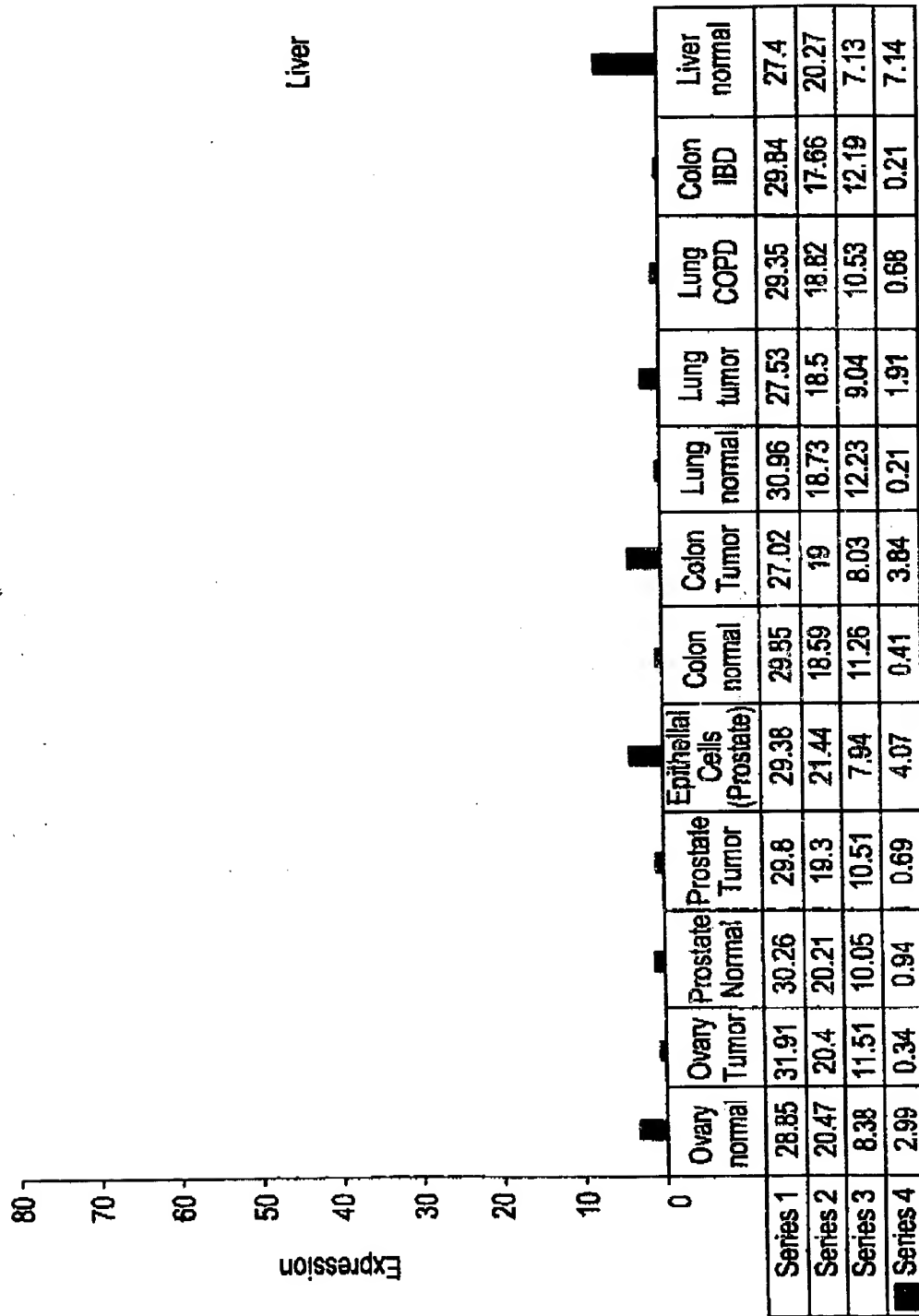
6177424214

T-652 P.49/54 F-983

27/32

Liver

FIG. 8C



May-17-02 09:37pm

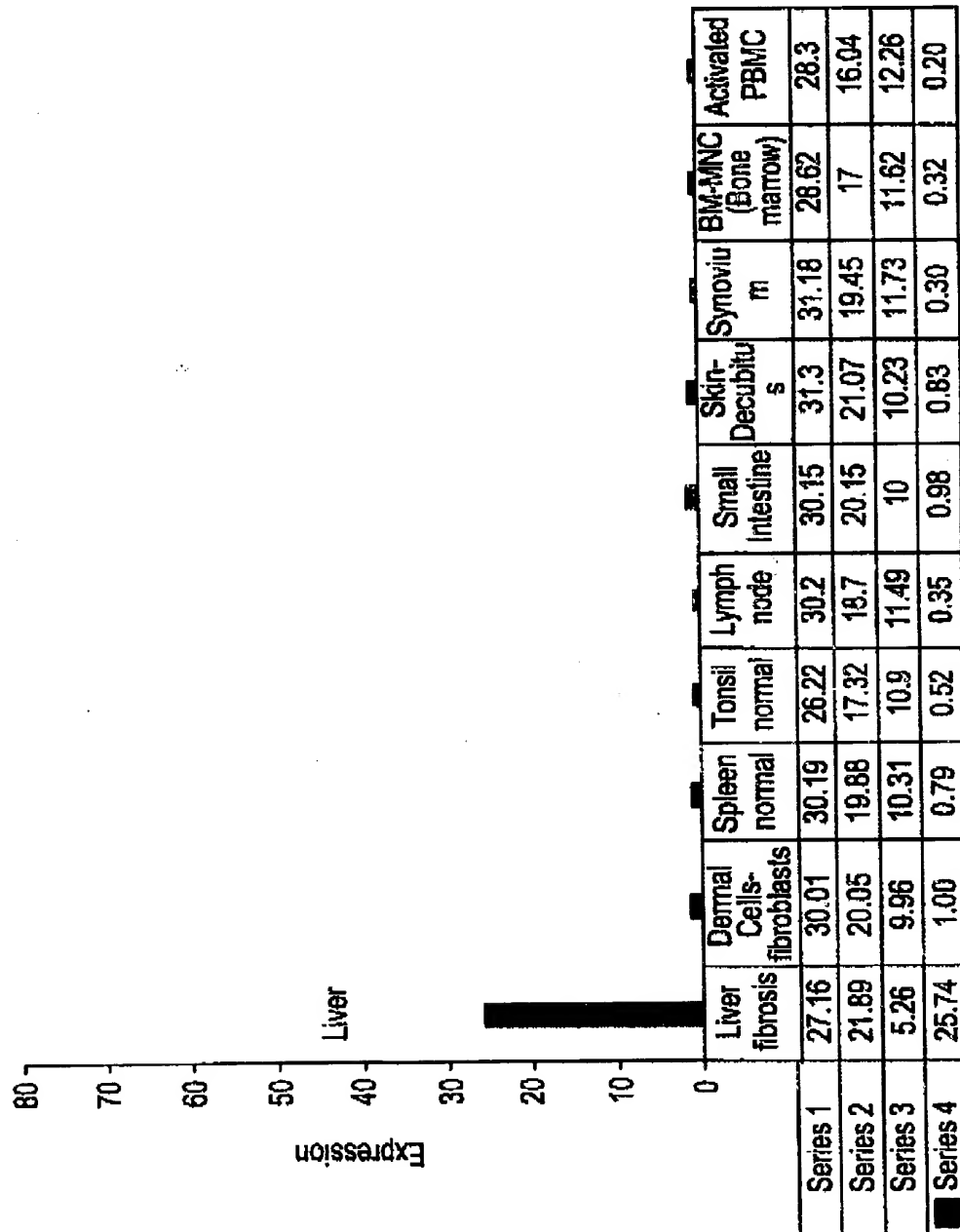
From-LAH1 COCKFIELD, LLP

6177424214

T-652 P.50/54 F-983

28/32

FIG. 8D



May-17-02 09:37pm

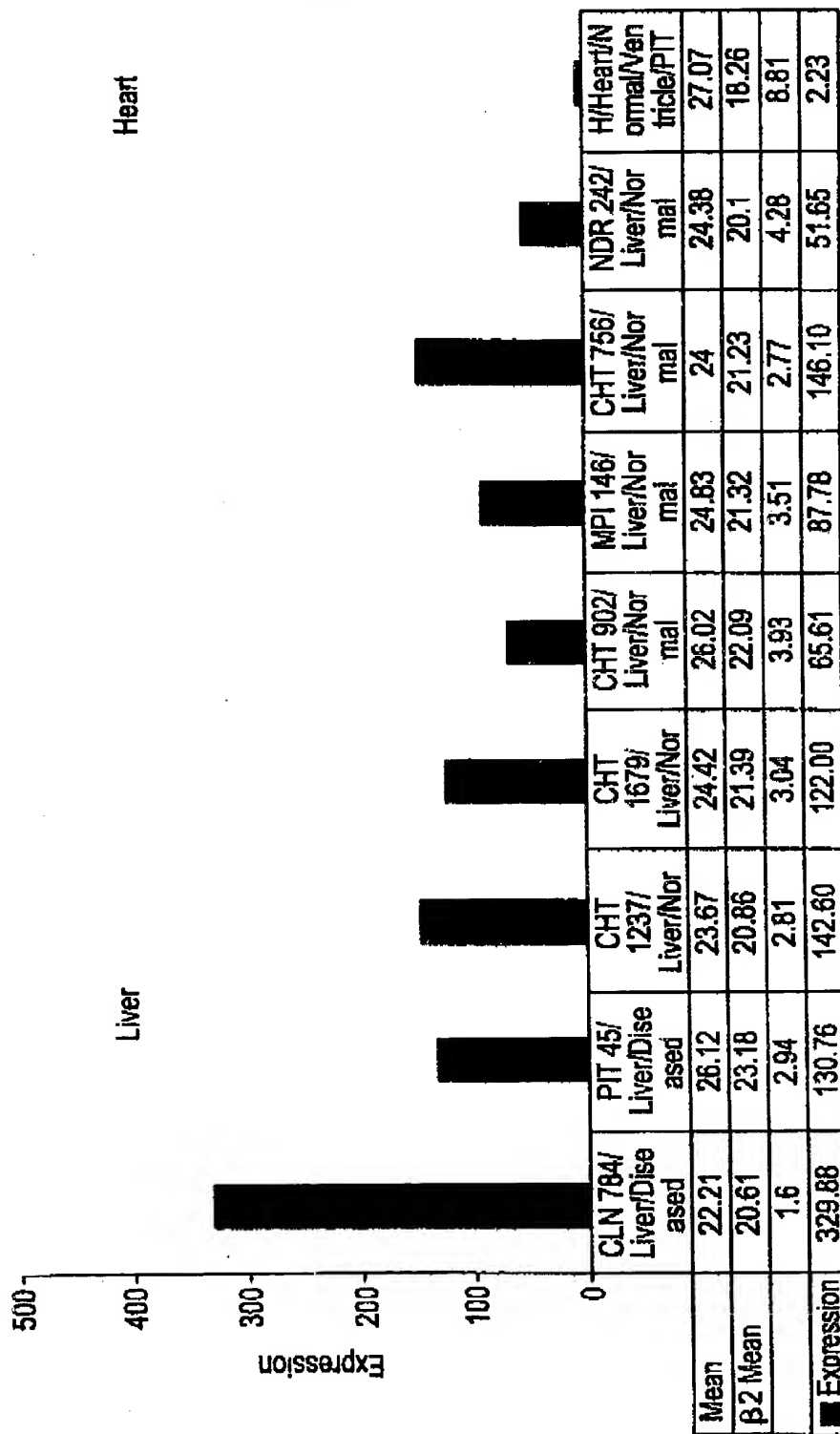
From-LAH1 ROCKFIELD, LLP

6177424214

T-652 P.51/54 F-983

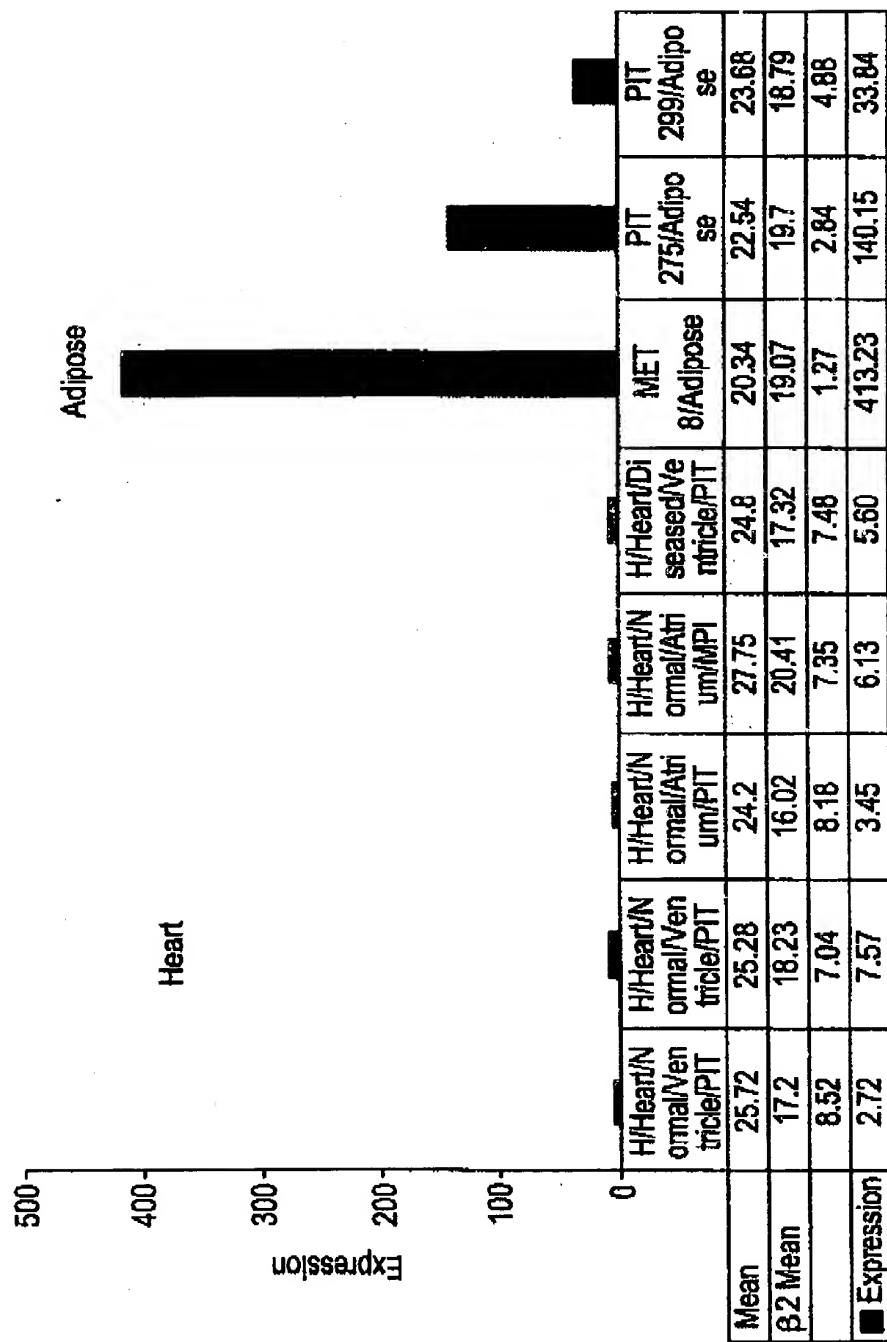
29/32

FIG. 9A



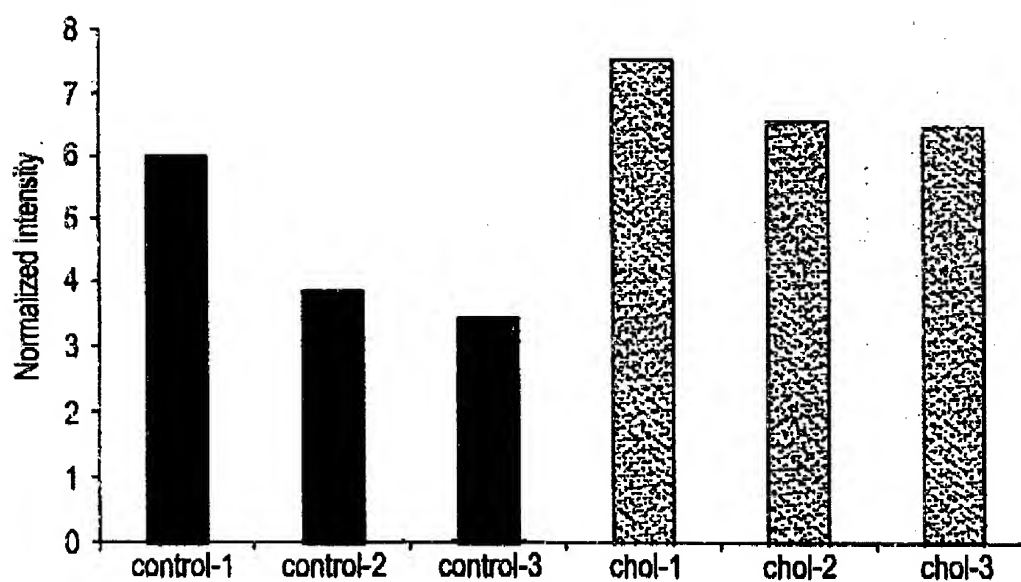
30/32

FIG. 9B



31/32

FIG. 10A



32/32

FIG. 10B

